PCI

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5:		(11	1) International Publication Number: WO 91/18926
C97K 15/04, C12N 15/31	A1	1,,,	3) International Publication Date: 12 December 1991 (12.12.91)
A61K 39/102 // C12Q 1/04		(43	5) International 1 noncomon Date:
C12Q 1/68, C12N 15/62	<u> </u>	Ц,	
(21) International Application Number: PCT/SI	E91/00	129	Published . With international search report.
(22) International Filing Date: 21 February 1991	(21.02.	91)	- "
(30) Priority data: 9001949-8 31 May 1990 (31.05.90)		SE	-
(71)(72) Applicant and Inventor: FORSGREN, Arne Sånekullavägen 33, S-217 74 Malmö (SE).	[SE/S	E];	
(74) Agent: AWAPATENT AB; Box 5117, S-200 (SE).	71 Mal	lmö	
(81) Designated States: AT (European patent), AU, pean patent), CA, CH (European patent), DE patent), DK (European patent), ES (European FI, FR (European patent), GB (European patent), IT (European patent), IP, pean patent), NL (European patent), NO, SE patent), US.	in pate atent), (LU (Et	nt), GR	
			WATERODULI IIC INCLUENZAE
(54) Title: PROTEIN D - AN IgD-BINDING PRO	TEIN	OF	HAEMOPHILOS INFLOENZAD
168 AAAAAAGGGGGGGGGGAAAAAGGGGGGGGGGGGGGGG	TETRALAMETO		178 ATTENTIONAL AND CONTRACTOR AND
119 ATAMANTTEACOGCACTOTAMOCACAMATACTISTCAMOCTAMATACAMOCTAMOCT	ACT HOUSE	= "	27 E28 CTTINGTHCANCIFICATITYANTHATANACTALLANGE STORES OF VALTY LONGISTER PROAFFINANCIAL LONGISTER STORES OF THE ST
223 TOTTENTION OF THE PROPERTY	ANTEROGRAPH ANTEROGRAPH ANTEROGRAPH	AT 21	67 CLASTOCKAPOCKATTONATTACTACLA TATIONAL PROPERTY AND CLASSIC CAMPACTURE AND CONTRACTOR AND CONT
188 ACCOMMUNICATION AND ANTICOMETRIC TOUTCOME TRACTION OF THE PROPERTY AND ANTICOMETRIC TOUTCOME TRACTION OF THE PROPERTY AND ANTICOMETRIC TOUTCOME TRACTION OF THE PROPERTY AND ANTICOMETRIC TOUTCOME TO THE PROPERTY AND ANTICOMETRIC TOUTCOMETRIC TOUTCOME TO THE PROPERTY AND ANTICOMETRIC TOUTCOMETRIC TOUTCOME TO THE PROPERTY AND ANTICOMETRIC TO THE PROPERTY AND ANTICOMETRIC TOUTCOME TO THE PROPERTY AND ANTI	CONTRACTOR CONTRACTOR	20 E7 3	143 Cleciniquiepreipedly Tysty Tellusity sanity singly the Pacipates Cleciniquiepreipedly Tysty Tellusity sanity singly the Pacipates
243 CACATACATTACACATTACACATTACACATACACACACA	riki iridas Pytlasiisi	ah (1998 CONCLUNICATION TO THAT THE CONTROL OF THE PROPERTY OF THE LOSS OF THE PROPERTY OF THE PRO
468 CATTIGCARTON-TRACTATOLITITIS CONTRACTA Application for the part of the par	CTTTTTAKKT	er '	1944 GTTAXTALCAMEATTEMACTCATARTETATION LLCG VOLCENTIALISM
464 Timerokroticomannekrociechcorekrotichterpannekrociechterpa	COUPLIANT OF THE	AFT .	577 1128 CENCATEVANTOTORIAGTECKFOTT TACACOTORI TALBANTOTORIAGTTT 1127 Alcoladyraanska (qiwo lalacyretyrime valkeyly saapala laceyredustes
. SIS ATOMETTALETTAAMMILITEALETTAMAMILITEALETTAMAMILIAMAA. Lalajamamiliamiliamiliamiliamiliamiliamil			1113 TYCHCHOTANYCAMITTANTHITOCTATTCANTANTCAGCAGAGATTA 1247 PROTECTION TO THE PROTECTION OF THE PROTECT
148 CATOCOLANDACOMOTENTETALACOTTECCATATOCALETTOCAL AMPLIQUIMENTALITY PROMOTENT PROPERTY I			947 1248 TTANTESTTECCHARACTOCCCCCAATCTTAAACCAATTAATTAATTAATCCCCCA 1287 PARTICASPROFTMANDTREELYTSIGIG Processignily 21 sapatind
449 ATTUCKBURTUMATUMATUMATUKANTATUKACAT PAGAMA 1441 ATTUCKBURTUMATUMATUMATUKANTATUKACAT PAGAMA 1441 ATTUCKBURTUMATUMATUMATUKANTATUKACAT PAGAMATUMATUMATUMATUMATUMATUMATUMATUMATUMAT			107 . 1308 CALCOSTOCIAMOSTACCACCOTTANTACCTICATATACCTICALACTAMATC (315)
140 AACTROONTENTCAEALTCAMCACTTOTTCCACCAC typicilyllatychiciallolpalabestymenialiae			747
·	_		
(57) Abstract			the anothin
A novel surface exposed protein of Haemon	hilus i	oflue	enzae or related Haemophilus species is described. The protein parent molecular weight of 42,000. Protein D can be detected in

A novel surface exposed protein of Haemophilus influenzae or related Haemophilus species is described. Into protein D is an Ig receptor for human IgD and has an apparent molecular weight of 42,000. Protein D can be detected in all of 116 encapsulated and non-encapsulated isolates of H. influenzae studied. The protein from all strains shows in addition to the same apparent molecular weight immunogenic similarities since protein D from all strains interacts with three different mouse monoclonal antibodies and monoclonal human IgD. A method for purification of protein D is described. Cloning of the protein D gene from H. influenzae in E. coli is described as well as the nucleotide sequence and the deduced amino acid sequence.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

DR CZ CZ CZ CZ CZ CZ CZ CZ CZ CZ CZ CZ CZ	andglafi Gertlina Faso	MC FIN KB KB LL LL LL CGB CGB CGB CGB CGB CGB CGB CGB CGB CGB	France Gabon United Kingdom Outmen Greece Hungary Isayan San Secondic People's Republic Mepublic of Korea of Korea San Laetenstein Sal Lanta Sal Lanta Handa	MINI MINI MINI MINI MINI MINI MINI MINI	Memorpolia Melawi Melawi Melawi Merakerands Memoras Potand Somenia Sweden Sweden Chnat Chnat Togo	
			Gabon			

1

PROTEIN D - AN IGD-BINDING PROTEIN OF HAEMOPHILUS INFLUENZAE

The present invention is related to a surface exposed protein named protein D which is conserved in many strains of Haemophilus influenzae or related Haemophilus species. Protein D is an Ig receptor for human IgD.

Several immunoglobulin (Ig) binding bacterial cell wall proteins have been isolated and/or cloned during the last two decades. The best characterized of these are protein A of Staphylococcus aureus and protein G of group G beta-hemolytic streptococci. The classical Fc-binding capacity of protein A involves IgG from humans and several mammalian species but the binding is restricted to human IgG subclasses 1, 2 and 4. Also other human classes of Ig (G, A, M, E) have been shown to bind to protein A, a reactivity that has been designed the alternative Ig binding which is mediated by Fab structures and characterized by a variable occurrence in the different Ig classes.

20 Protein G of group G streptococci binds all human IgG subclasses and has also a wider binding spectrum for animal IgG than protein A. On the IgG molecule the Fc part is mainly responsible for the interaction with protein G although a low degree of interaction was also recorded for 25 Fab fragments. IgM, IgA and IgD, however, show no binding to protein G. Both protein A and protein G have acquired many applications for immunoglobulin separation and detection. (EP 0 200 909, EP 0 131 142, WO 87/05631, US 3,800,798, US 3,995,018.)

Certain strains of group A streptococci are also known to produce an IgG-binding protein which has been purified or cloned. The Ig-binding protein from group A streptococci is relatively specific for human IgG. Information about bacterial molecules that selectively bind IgA and IgM is more limited. However, IgA-binding proteins have been isolated from both group A and group B streptococci, two frequent human pathogens. The IgA receptor of

30

group A streptococct has been named protein Arp. Certain strains of the anaerobic bacterium Clostridium perfringens preferentially bind igm but also igh and igG. This binding beferentially bind igm but also igh and igG. This binding bacterial protein, protein (protein P). Recently a ties for L-chains was isolated from Peptococcus magnus. Protein L has been shown to bind igG, igh and igM from bacteria, ig receptors have been reported among veterinary pathogens. Brucella abortus binds bovine igM and Taylorella equigentialis, a venereal pathogen of horses, binds equigentialis, a venereal pathogen of horses, binds equine igG. Recently Haemophilus somnus was reported to bind bovine igG. Recently Haemophilus somnus was reported to

(Branhamella) catarrhalis were shown to have a high binding capacity for human IgD (Forsgren A. and Grubb A, J. Immunol. 122:1468, 1979).

A decade ago Haemophilus influenzae and Moraxella

Protein D is not identical with any previously described tmmune system through an interaction with B-lymphocytes. proteins and polysaccharides) in the stimulation of the stone and in combination with other molecules (for example used for IgG. Protein D could also be a valuable tool wel tn which protein A and protein G previously have been separation and detection of IgD in a way similar to the Thus, protein D could be an important tool for studies, with IgD and not with other human immunoglobulin classes. Ig-binding proteins. Protein D was found only to interact found to be different compared with previously isolated properties of this molecule, named protein D, which were richia coli. In addition it describes the Ig-binding IgD-binding protein gene of the H. influenzae in Eschethe cloning, expression and nucleotide sequence of the ponsible for the interaction with IgD. It also describes and purification of a H. influenzae surface protein res-The present invention describes the solubilization

protein from H. influenzae.

H. influenzae is a common human parasite and pathogen which colonizes the mucosa of the upper respiratory tract and causes disease by local spread or invasion. An important distinguishing feature between H. influenzae isolates 5 is whether or not they are encapsulated. Encapsulated H. influenzae type-b is a primary cause of bacterial meningitis and other invasive infections in children under 4 years of age in Europe and the United States. Non-encapsulated (non-typable) H. influenzae rarely cause invasive 10 infection in healthy children and adults but are a frequent cause of otitis media in children and have been implicated as a cause of sinusitis in both adults and children. H. influenzae are also commonly isolated in purulent secretions of patients with cystic fibrosis and 15 chronic bronchitis and have recently been recognized as an important cause of pneumonia.

A vaccine composed of purified type b capsular polysaccharide has proven effective against H. influenzae type
b disease in children of 2 to 5 years of age. However,
20 since children under two years of age respond poorly to
this vaccine, conjugate vaccines with enhanced immunogenicity have been developed by covalently bonding the capsular polysaccharide to certain proteins. However, the
polysaccharide vaccines, non-conjugated and conjugated,
25 are of no value against nontypable H. influenzae disease.
Hence, other cell surface components and in particular
outer membrane proteins (OMPs) have been looked at as
potential vaccine candidates both against type b and
nontypable H. influenzae. (EP 0 281 673, EP 0 320 289.)

The outer membrane of H. influenzae is typical of gram-negative bacteria and consists of phospholipids, lipopolysaccharide (LPS), and about 24 proteins. Four different Haemophilus OMPs have been shown to be targets for antibodies protective against experimental Haemophilus disease. These include the Pl heat-modifiable major outer membrane protein, the P2 porin protein, the P6 lipoprotein and a surface protein with an apparent molecular weight of

30

MO 31\18379 bCL\ZE31\00173

98,000 (98 K-protein). Of these at least antibodies to P2 have been shown not to protect against challenge with heterologous Haemophilus strains. (Loeb, M. R. Infect. Immun. 55:2612, 1987; Munson Jr, R. S. et al J. Clin. Invest. 72:677, 1983; Munson Jr, R. S. and Granoff, D. M. Infect. Immun. 49:544, 1985 and Kimura, A. et al, Infect. Immun. 194:495, 1985).

there are marked differences in OMP composition among strains (See e.g. Murphy et al. "A subtyping system for nontypable Haemophilus influenzae based on outer membrane proteins" J Infect Dis 147:838, 1983; Barenkamp et al. "Outer membrane protein and biotype analysis of pathogenic nontypable Haemophilus influenzae" Infect Immun 30:709, "Outer membrane protein and biotype analysis of pathogenic nontypable Haemophilus influenzae" Infect Immun 30:709, "Outer membrane protein and pathogenic nontypable Haemophilus influenzae ha

Analysis of nontypable H. influenzae has shown that

If a surface exposed antigen (immunogen) which is conserved in all strains of H. influenzae could be found it would be an important tool in developing a method of identifying H. influenzae. The present invention shows that protein b with an identical apparent molecular shows that protein b with an identical apparent monoclonal weight (42,000), reacting with three different monoclonal antibodies and human igb, was found in all li6 H. influence of the constant of the constant

Thus, according to the invention there is provided a surface exposed protein, which is conserved in many strains of Haemophilus influencae or related Haemophilus a capacity of binding human 1gb. The invention also comprises naturally occurring or artificially modified one prises naturally occurring or artificially modified one—

variants of said protein, and also immunogenic or 1gbvariants of said protein, and variants. The pro-

H. haemolyticus and H. aegypticus.

variants of said protein, and also immunogenic or IgD-binding portions of said protein and variants. The protein is named protein D and has the amino acid sequence
depicted in Fig. 9.

There is also provided a plasmid or phage containing _ a genetic code for protein D or the above defined variants or portions.

Further there is provided a non-human host containing
the above plasmid or phage and capable of producing said
protein or variants, or said portions thereof. The host is
chosen among bacteria, yeasts or plants. A presently preferred host is E. coli.

In a further aspect the invention provides for a DNA segment comprising a DNA sequence which codes for protein D, or said variants thereof, or for said portions. The DNA sequence is shown in Fig. 9.

In yet another aspect, the invention provides for a recombinant DNA molecule containing a nucleotide sequence coding for protein D, or said variants or portions, which nucleotide sequence could be fused to another gene.

A plasmid or a phage containing the fused nucleotide defined above could also be constructed.

Further such a plasmid or phage could be inserted in 20 a non-human host, such as bacteria, yeasts or plants. At present, E. coli is the preferred host.

The invention also comprises a fusion protein or polypeptide in which protein D, or said variants or portions, could be combined with another protein by the use of a recombinant DNA molecule, defined above.

Furthermore, a fusion product in which protein D, or said variants or portions, is covalently or by any other means bound to a protein, carbohydrate or matrix (such as gold, "Sephadex" particles, polymeric surfaces) could be constructed.

The invention also comprises a vaccine containing protein D, or said variants or portions. Other forms of vaccines contain the same protein D or variants or portions, combined with another vaccine, or combined with an immunogenic portion of another molecule.

30

modified variants thereof. . of protein D, or of naturally occurring or artificially producing a monoclonal antibody to an immunogenic portion There is also provided a hybridoma cell capable of

the antibody in the presence of an indicator. byijne sbecies in a sambje by contacting said sample with the presence of Haemophilus influenzae or related Haemothereof. This antibody is used in a method of detecting naturally ooccurring or artificially modified variants is specific to an immunogenic portion of protein D or of Further there is provided a purified antibody which

occurring or artificially modified variants thereof, or bytine sbecies in a sample by contacting said sample with the presence of Haemophilus influenzae or related Haemo-The invention also comprises a method of detecting

or variants. for an immunogenic or IgD-binding portion of said protein nucleic acids which code for protein D, or for naturally a DNA probe or primer constructed to correspond to the

Finally, the invention comprises a method of sepathe protein may be labelled or bound to a matrix. in a method of detecting IgD. In such a detecting method Protein D, or said variants or portions, is also used

optionally bound to a matrix. rating IgD using protein D, or said variants or portions,

MATERIALS AND METHODS

Bacteria

20

ting l2 species related to H. influenzae were obtained 116 H. influenzae strains representing serotypes a-f

.A.2.U from different laboratories in Denmark, Sweden and the and nontypable and in addition bacterial strains represen-

Culture conditions

philus strains in an atmosphere containing 5% ${\rm CO}_2$. 30 microserophilic atmosphere at 37°C and all other Haemolus were grown on chocolate agar. H. ducreyi were grown in All strains of Haemophilus, Ekinella and Acinobacilisolates of H. influenzae were also grown overnight at 37°C in brain-heart infusion broth (Difco Lab., Inc. Detroit, Mi.) supplemented with nicotinamide adenine dinucleotide and hemin (Sigma Chemical Co. St Louis, Mo.), each at 10 µg/ml.

Immunoglobulins and proteins

IgD myeloma proteins from four different patients were purified as described (Forsgren, A. and Grubb, A., J. Immunol. 122:1468, 1979). Eight different human IgG myeloma proteins representing all four subclasses and both L-chain types, three different IgM myeloma proteins and one IgA myeloma protein were isolated and purified according to standard methods. Human polyclonal IgG, serum albumin and plasminogen were purchased from Kabi Vitrum 15 AB, Stockholm, Sweden, and human IgE was adapted from Pharmacia IgE RIACT kit (Pharmacia Diagnostic AB, Uppsala, Sweden). Bovine serum albumin, human and bovine fibrinogen and human transferrin were purchased or obtained as a gift.

20 125 I-IgD binding assay

The binding assay was carried out in plastic tubes. Briefly 4×10^8 bacterial cells in a volume of 100 µl phosphate buffered saline (PBS) with the addition of 5% human serum albumine (HSA) were mixed with 100 µl of 125 I-IgD in the same buffer (radioactivity was adjusted to 7-8×10⁴ cpm, i.e approx. 40 ng). After 0.5 h incubation at 37°C, 2 ml of ice-cold PBS (containing 0.1% Tween 20) was added to the tubes.

The suspension was centrifugated at 4,599xg for 15 min and the supernatant was aspirated. Radioactivity retained in the bacterial pellet was measured in a gamma counter (LKB Wallac Clingamma 1271, Turku, Finland). Residual radioactivity from incubation mixtures containing no bacteria, i.e. background, was 2.5 percent. Samples were always tested in triplicates and each experiment was repeated at least twice, unless otherwise stated.

Monoclonal antibodies

phoresis was performed at room temperature using PROTEIN 10% glycerol, and 0.03% (w/v) bromphenol blue. Electro-Tris hydrochloride (pH 6.8), 2% (W/V) SDS, 1% (VV) β -ME, by 5-min boiling in sample buffer consisting of 0.06M of influenzae and related bacterial species were pretreated ration of 11%. Samples of crude Sarcosyl extracts of H. (FEBS Lett 58:254, 1975) using a total acrylemide concentand run according to the procedure of Lugtenberg et al., SDS-PAGE was, using a modified Laemmii gel, prepared SDS-PAGE and detection of protein D on membranes and 90% fetal bovine serum in liquid nitrogen. Tives were trozen in the presence of dimethyl sulfoxide selected for further growth in the same medium. All cell to protein D were obtained. Three of the hybridomas were fetal bovine serum. Totally 68 clones producing antibodies expanded by cultivation in RPMI medium containing 10% producing the highest titers of antibodies were cloned and -linked immunosorbent assay (ELISA), and the hybrids production of antibodies against protein D in an enzyme-14 days (mean 12 days) the hybridomas were tested for the Scheidegger SJ J Immunol Methods 35:1, 1980). After 10 to duction of monoclonal antibodies (De St Groth SF, was excised and spleen cells were prepared for the pro-150 µl PBS. One day after the last injection, the spleen poosted by an intravenous injection of protein D (2 µg) in sorbent assay (ELISA). The best responding mouse was for anti-protein D activity in an enzyme-linked immunowere bled from the tails, serum was separated and tested adjuvant (300 µl) 3 and 7 weeks later. In week 9 the mice injections of protein D (15 µg) in Freund's incomplete saljuvant (300 µl) followed by two intraperitoneal purified protein D (25 µg/50 µl) in Freund's complete immunized by an intraperitoneal injection of 25 µg Inbred female BALB/c mice (age 8 to 14 weeks) were

II vertical slab electrophoresis cells (Bio-Rad Laboratories, Richmond, CA) at 40 mA per gel constant current. Staining of proteins in gels was done with comassie brilliant blue in a mixture of methanol, acetic acid and water essentially as described by Weber and Osborn (J. Biol. Chem. 244:4406, 1969). Protein bands were also transferred to nitrocellulose membranes (Sartorius, West Germany) by electrophoretic transfer from SDS-polyacrylamide gels. Electrophoretic transfer was carried out in a Trans-Blot Cell (Bio-Rad) at 50 V for 90 min. The electrode buffer was 0.025M Tris, pH 8.3, 0.192M glycine, and 20% methanol. The membranes were then washed for 1 h at room temperature in 1.5% ovalbumin-Tris balanced saline (OA-TBS), pH 7.4, to saturate additional binding sites.

After several washings with Tris balanced saline (TBS), the membranes were incubated overnight at room 15 temperature in 1% OA-TBS buffer containing IgD (20 µg/ml). to detect IgD-binding bands, then washed twice with TBS... The membranes were then incubated with peroxidase conjugated goat anti-human IgD (Fc) (Nordic Immunology, Tiiburg, The Netherlands) for 1-2 hrs at room temperature; after several washings with Tween-TBS the membranes were developed with 4-chloro-1-napthol and hydrogen peroxide. Protein D was also identified using anti-protein D mouse monoclonal antibodies 16Clo, 20G6 and 19B4 at 1:50 dilution in 1% OA-TBS. Protein 1 and 2 of H. influenzae 25 were identified using anti-P2 mouse monoclonal 9F5 (Dr. Eric J. Hansen, Dallas, Texas, USA) at a 1:1000 dilution and rabbit anti-Pl serum (Dr. Robert S. Munson, St. Louis, Mo, USA) at a 1:200 dilution. Solubilization and purification of protein D from H. 30 influenzae

Briefly 3 g of bacteria were suspended in 10 ml of 10 mM HEPES Tris buffer (pH 7.4) containing 0.01M EDTA and sonicated three times in a sonifier (MSE) for 1 min while cooling in an ice bath. Following sonication Sarcosyl

35 (Sodium Lauryl Sarcosinate) was added to a final concentration of 1% (w/v). The suspensions were incubated at room temperature for 1 h using a shaker and then sonicated

MO 31/18379 FCL/SE31/00173

OT

again 2xl min on ice and reincubated at room temperature for 30 min. After centrifugation at 12,000 g for 15 min at 4° C the supernatant was harvested and recentrifugated at 4° C the supernatant was harvested and recentrifugated at

dialysis was performed. Finally the supernatant was concentrated and extensive stter 4 hrs at 4°C centrifugation was performed as above. after the potassium content was adjusted to 60 mM and pitate was removed by centrifugation at 12,000 g. Thereadded and after incubation at 4°C overnight the SDS-preci-Potassium phosphate in a final concentration of 20 mM was trom Susukt and Terrada (Anal. Biochem. 172:259, 1988). precipitation in potassium phosphate buffer using a method was removed from the protein containing solution by Igb-binding molecules (protein D) was performed and SDS identified and cut out. Electrophoretic elution of the (Western blot) the appropriate band in the gel could be compartson with the IgD-binding band on the membrane 2DS-PAGE and detection of protein D on membranes). By conjudated gost anti-human IgD as described above (see detected by Western blot assay using IgD and peroxidase was transferred to membranes and the IgD-binding band was electrophoresis narrow gel strips were cut out, protein 772 as described above were applied to SDS-PAGE. After Sarcosylextracts prepared of H. influenzae, strain NT

Proteins were applied to nitrocellulose membranes (Schleicher & Schuell, Dessel, West Germany) manually by using a dot blot apparatus (Schleicher & Schuell). After temperature in 1% OA-TBS containing 125 I-labeled protein probe (5 to 15 IO), washed four times with TBS containing 0.02% Tween-20, air dried, and sutoradiographed at -70°C by using Kodak CEA.C X-ray films and Kodak X-Omat at reqular intensifying screen (Eastman Kodak, Rochester, regular intensifying screen (Eastman Kodak, Rochester,

.(УИ

Dot blot assay

Amino acid sequence analysis

Automated amino acid sequence analysis was performed with an Applied Biosystems 470A gas-liquid solid phase sequenator (A) with online detection of the released amino acid phenylthiohydantoin derivatives by Applied Biosystems Model 120A PTH Analyzer.

Bacterial strains, plasmids, bacteriophages and media used for cloning of protein D

H. influenzae, nontypable strain 772, biotype 2, was isolated from a nasopharyngeal swab at the Department of Medical Microbiology, Malmö General Hospital, University of Lund, Sweden. E. coli JM83 were used as recipient for plasmids pUC18 and pUC19 and derivatives thereof. E. coli JM101 and JM103 were used as hosts for M13mpl8 and mp19 bacteriophages. H. influenzae was cultured in brain-heart infusion broth (Difco Lab., Inc. Detroit, Mi.) supplemented with NAD (nicotine adenine dinucleotide) and hemin (Sigma Chemical Co., St Louis, Mo.), each at 10 µg/ml. E. coli strains were grown in L broth or 2xYT media. L agar and 2xYT agar contained in addition 1.5 g of agar per litre. L broth and L agar were, when so indicated, supplemented with ampicillin (Sigma) at 100 µg/ml.

Chromosomal DNA was prepared from H. influenzae

25 strain 772 by using a modification of the method of Berns and Thomas (J Mol. Biol. 11:476, 1965). After the phenol:chloroform:isoamylalcohol (25:24:1) extraction step the DNA was ethanol precipitated. The DNA was dissolved in 0.1xSSC (1xSSC:0.15 M NaCl and 0.015 M sodium citrate) and

30 RNase treated for 2 h at 37°C. The RNase was removed with two chloroform:isoamylalcohol (24:1) extractions. The DNA was banded in a CsCl-ethidium bromide equilibrium gradient.

Plasmid DNA and the replicative form of phage M13
35 from E. coli JM101 were obtained by the alkaline lysis procedure followed by further purification in a CsCl-ethidium bromide gradient. In some cases plasmid DNA was

ΖŢ

prepared using a Quiagen plasmid DNA kit (Diagen GmbH Düsseldorf, FRG).

Single-stranded (ss) DNA from phage Ml3 clones was prepared from single plaques (Messing, J. Meth. Enzymol

Molecular cloning of the protein D gene
A H. influensee genomic library was constructed

starting from 40 µg of H. influenzae strain 772 DNA which was partially digested with 1.2 units Sau3A for 1 h at 10 37°C. The cleaved DNA was fractionated on a sucrose gradient (Clark-Curtiss, J. E. et al., J. Bacteriol. 161:1093, 1985). Fractions containing DNA fragments of

gradient (Clark-Curtiss, J. E. et al., J. Bacteriol.
161:1093, 1985). Fractions containing DNA fragments of
appropriate sizes (2-7 kilobasepairs (kbp)) were pooled
and the DNA was ligated to dephosphorylated BamHI digested
5 pUCl8 under standard conditions (Maniatis, T. et al.,
Molecular cloning: A laboratory manual, 1982). The

Molecular cloning: A laboratory manual, 1982). The Ligation mixture was transformed into component E. coli TM83 by high voltage electroporation with a Gene Pulser TM/Pulse controller apparatus, both from Bio-Rad Lab.

(Richmond, CA). The bacteria were plated onto L agar

(Richmond, CA). The bacteria were plated onto L agar

-3-indolyl-6-D-galactopyranoside). Colony immunoassay

For colony immunoblotting, E. coli transferred to nitro-

cellulose filters (Sartorius GmbH, Göttingen, FRG) by covering the agar surfaces with dry filters. The plates were left for 15 min before the filters were removed and exposed to saturated chloroform vapour for 15 min. Resinnoubating the filters on the filters were blocked by froubating the filters in Tris balanced saline containing novalbumine for 30 min (TBS-ova; 50 mM Tris-HCL, 154 mM ovalbumine for 30 min (TBS-ova; 50 mM Tris-HCL, 154 mM incubated in turn with (1) culture supernatants containing from turn with (1) culture supernatants containing and an arm with (1) culture supernatants containing from the filters were some monoclonal antibodies (Wabs) directed against protein D at a dilution of 1:10 in TBS-ova, (ii) horseradish tein D at a dilution of 1:10 in TBS-ova, (ii) horseradish

peroxidase conjugated rabbit anti-mouse IgGs (DAKOPATTS

10

A/S, Glostrup, Denmark) in TBS-ova at a dilution of 1:2000 in TBS-ova, and (iii) 4-chloro-l-naphthol and ${\rm H_2O_2}$. The filters were washed 3xlO min in wash buffer (TBS-0.05% Tween 20) between each step. All incubations were done at room temperature.

Colonies were also checked for IgD binding by incubating other filters with purified human myeloma IgD:s, rabbit anti-human IgD (δ -chains) (DAKOPATTS), horseradish peroxidase conjugated goat anti-rabbit Ig:s (Bio-Rad Lab.) and 4-chloro-l-naphthol and ${\rm H_2O_2}$ as above.

Restriction endonuclease analysis and DNA manipulations

Plasmid and phage DNA were digested with restriction endonucleases according to the manufacturers' instructions (Boehringer Mannheim mbH, Mannheim, FRG, and Beckman Instruments, Inc., England). Restriction enzyme fragments for subcloning were visualised with low energy UV-light.

and excised from 0.7-1.2% agarose gels (Bio-Rad) containing 0.5% ethidium bromide. The DNA bands were extracted with a Geneclean TM kit (BIO 101 Inc., La Jolla, Ca.) as recommended by the supplier.

Ligations were performed with 14 DNA ligase (Boehringer Mannheim) under standard conditions (Maniatis et al., 1982). The ligation mixtures were used to transform competent E. coli cells.

pHIC348 for the sequencing procedure were produced by varying the time of exonuclease III digestion of KpnI-BamHI-opened plasmid DNA (Henikoff, S. Gene 28:351, 1984). For removal of the resulting single-stranded ends, mung bean nuclease was used. Both nucleases were obtained from Bethesda Research Laboratories Inc. (Gathersburg, Md.).

Protein D extraction from E. coli

Cells of E. coli expressing protein D were grown in L broth supplemented with ampicillin to early logarithmic phase and then subjected to osmotic shock. After removal of periplasmic fraction the cells were lysed with NaOH

(Russel, M. and Model, P., Cell 28:177, 1982) and the accetic acid.

Traction by centrifugation. The periplasmic and cytofraction by centrifugation. The membrane
fractic acid.

DNA sequencing and sequence manipulations

The nucleotide sequence was determined by direct plasmid sequencing (Chen, E. Y. and Seeburg, P. H. DNA 4:165, 1985) of subclones and deletion derivatives of d_1^{35} of subclones and deletion method with d_1^{35} slochemical Corp., Cleveland, Ohio) following the States Biochemical Corp., Cleveland, ohio) following the protocol provided by the supplier. Part of the sequencing protocol provided by the supplier. Part of the sequencing was done on single-stranded M13 DNA carrying inserts was done on single-stranded M13 DNA carrying inserts

Fult X-ray film. RESULTS Distribution of protein D in Haemophilus influenzae

A total of 116 H. influenzae strains obtained from

culture collections and freshly isolated from nasopharyngeal swabs were selected for 1gD-binding experiments.

Eleven of the strains were encapsulated representing serotypes a-f, and 105 strains were non-encapsulated (nontypable). These 105 strains belonged to biotype I (21 strains), biotype II (39 strains), biotype II (14 strains), biotype IV (2 strains) and biotype II (14 strains), biotype IV (2 strains) and biotype I (5

strains). Of the non-encapsulated strains 31 were not blotyped (NBT) but tested for 1gD binding.

Approximately 4x10⁸ cfu of H. influenzae bacteria grown on chocolate agar were mixed and incubated with 40 ng of radiolabeled human myeloma 1gD. Thereafter a

40 ng of radiolabeled human myeloma 1gD. Thereafter a larger volume (2 ml) of PBS containing Tween 20 was added, bacteria were spun down and radioactivity of pellets was measured. All H. influences isolates bound 1gD to a high degree (38-74%) (Fig. 1). There was no difference in IgD-binding capacity between different serotypes (a-1) of encapsulated H. influences. Nor was there any difference

between different biotypes of non-encapsulated strains.

30 strains representing different sero- and biotypes were also grown in brain-heart infusion broth. When those bacteria grown in liquid medium were compared with the same bacteria grown on chocolate agar, no difference in IgD-binding capacity could be detected.

Protein D was solubilized from all 116 H. influenzae strains by sonication and Sarcosyl extraction. Subsequently the extracts containing protein D were subjected to SDS-PAGE. Proteins were stained or electroblotted onto nitrocellulose membranes and probed with human IgD myeloma protein and three different mouse monoclonal antibodies recognizing protein D. Many protein bands could be detected in all SDS-gels but electrophoresis of extracts from all H. influenzae isolates gave a protein band with an apparent molecular weight of 42,000 (42 kilodaltons). IgD and also all three anti-protein D monoclonal antibodies (16ClO, 20G6 and 19B4) bound to the same band after electrophoresis of all extracts and subsequent transfer to membranes and blotting.

Bacterial strains of 12 different species taxonomically related to H. influenzae (H. ducreyi, H. paraphrophilus, H. parasuis, H. parainfluenzae, H. haemolyticus, H. parahaemolyticus, H. aphrophilus, H. segnis, H. aegypticus, H. haemoglobinophilus, E. corrodens, A. actinomycetemcomitans) were tested for their capacity to bind 125 I labeled human IgD. In addition crude Sacrosyl extracts from the same bacteria were tested by Western blot analysis with IgD and the three anti-protein D monoclonal antibodies (MAbs 16C10, 20G6, 19B4).

Of all twelve species tested, only H. haemolyticus (5/5 strains) and H. aegypticus (2/2 strains) bound radiolabeled IgD, 21-28% and 41-48%, respectively, in the direct binding assay (Fig. 2). In Western blot analysis IgD and all three monoclonal antibodies detected a single band with an apparent molecular weight of 42,000 (42 kilodaltons).

Three different strains of H. influenzae (two non-02 Solubilization of protein D able by IgD or the three monoclonal antibodies. the same bacteria did not reveal any protein band detectigh in the direct binding assay and Sarcosyl extracts from H. haemoglobinophilus (1 strain) did not bind radiolabeled H. parahaemolyticus (2 strains), H. sengius (2 strains), bodies. Two strains of H.ducreyi, H. parasuis (2 strains), band was detected with the two other monoclonal antiextract of one of the strains, a single 42 kilodaltons daltons) with MAb 16ClO in all three strains. In an revealed a single high molecular weight band (90 kilo-Western blot analysis of three strains of E. corrodens lysis showing a single 42 kilodaltons protein band. three of the monoclonal antibodies in Western blot anaever, extracts of all these strains reacted with two or assay or reacted with IgD in Western blot analysis. Howcomitans bound radiolabeled IgD in the direct binding parainfluenzae, 8 H. aphrophilus, and 3 A. actinomycetem-Mone of the 6 strains of H. paraphrophilus, 11 H.

typable strains, 772 and 3198 and one type B, Minn A.)

were grown overnight in broth. Initially attempts were
made to solubilize protein D according to a well

established method for isolation of H. influenzae outer
membrane proteins by sonication, removal of the cell
debris by centrifugation and extraction of the supernatant
vith Sarcosyl followed by ultracentrifugation (Barenkamp
SJ and Munson RS J Infect Dis 143:668, 1981). The pellets
(cell debris) (d) and supernatants (s) after sonication as
syl-treatment and ultracentrifugation were subjected to
syl-treatment and ultracentrifugation were subjected to
syl-treatment and ultracentrifugation of the sonication as
protein followed by incubation with peroxidase conjugated
protein followed by incubation with peroxidase conjugated
anti-human igD-antibodies and substrate. As shown in

Fig. 3 the sonication procedure solubilized proteins including protein D effectively. However, IgD-binding

10

molecules (protein D) could also be detected in the cell debris, i.e. were not solubilized by sonication. The yield of IgD-binding molecules in the supernatant varied between different experiments. Fig. 3 also shows that protein D 5 mostly could be detected in the Sarcosyl soluble supernatant after ultracentrifugation. In contrast previously described outer membrane proteins of H. influenzae (protein 1 to 6) are readily solubilized by sonication and are considered Sarcosyl insoluble.

To improve the yield of protein D several extraction methods were tried. In subsequent experiments the bacterial cells were sonicated and the whole cell suspension sonicated and extracted in different detergents (Sarcosyl, NP-40, Triton X-100 and Tween 80). The cell debris was 15 removed by centrifugation (12,000 g) and the supernatant. ultracentrifuged. The thus obtained cell debris (d), supernatants (s) and pellets (p) were analysed by SDS--PAGE, electroblotting onto membranes and subsequent probing with IgD. As shown in Fig. 4 Sarcosyl treatment effectively solubilized protein D leaving little left in . the cell debris and pellet. NP-40, Triton X-100 and Tween--80 solubilized protein D less effectively.

Attempts were also made to solubilize protein D from the bacteria with lysozyme and different proteolytic enzymes (papain, pepsin and trypsin) at different concentrations. Of the enzymes only lysozyme solubilized protein D (Fig. 4).

Purification of protein D

Protein D was solubilized by Sarcosyl extraction of 30 whole bacteria as described above and purification was performed by SDS-PAGE of the supernatant after ultracentrifugation. After electrophoresis narrow gel strips were cut out, proteins were transferred to membranes and the IgD-binding band (protein D) was detected by Western blot assay. Gel slices containing a protein band corresponding to the IgD-binding molecules were cut out from the gel and solubilized by electronic elution. At reelectrophoresis

the purified protein, protein D (D), migrated as a single band (42 kilodaltons) (Fig. 5) without discernible breakdown products.

To confirm that protein D was not identical with the previously described outer membrane proteins 1 or 2 with molecular weights of 49 and 39 kilodaltons, respectively, of whole H. influencae bacteria were subjected to SDS-PAGE, transferred to Immobilon filters and blotted with antibodies to protein 1 and protein 2 and also with human antibodies to protein 1 and protein 2 and also with human light and protein 2 and size with human antibodies to protein 1 and protein 2 and also with human antibodies to protein 1 and protein 2 and size with human antibodies to protein 2 and size with human antibodies are seen in Fig. 5 protein 2 and size difference with human antibodies are seen in Fig. 5 protein 2 and size difference with human antibodies are seen in Fig. 5 protein 2 and size difference with human antibodies are seen in Fig. 5 protein 2 and size difference with human antibodies are seen in Fig. 5 protein 2 and size difference with human antibodies are seen in Fig. 5 protein 2 and size difference with human antibodies are seen in Fig. 5 protein 2 and size difference with human antibodies are seen in Fig. 5 protein 2 and size difference with human antibodies are seen in Fig. 5 protein 2 and size difference with human antibodies are seen in Fig. 5 protein 2 and size difference with human antibodies are seen in Fig. 5 protein 2 and size difference with human antibodies are seen in Fig. 6 and size difference with human antibodies are seen in Fig. 6 and size difference with human antibodies are seen in Fig. 6 and size difference with human antibodies are seen in Fig. 6 and size difference with human antibodies are seen in Fig. 6 and size difference with human antibodies are seen in Fig. 6 and size difference with human antibodies and size difference with human and size difference with human antibodies and size difference with human and size difference with human antibodies and size difference with human and size difference with

IgD. As can be seen in Fig. 5 protein D migrates differently from protein 1 and protein D

The interaction of protein D with human IgD was

further verified in gel filtration experiments where of the two protein D was eluted together with IgD when a mixture of the two protein D run alone on the same column was eluted 60). Protein D run alone on the same column was eluted 60). Protein D run alone on the same column was eluted 60). Protein D run alone on the same column was eluted 60). Protein D run alone on the same column of Fig. 7.

42 kilodaltons for protein D was also studied in different dot blot experiments to further examine the binding speci-

ficity of the molecule. Fig. 7 shows that protein D

ficity of the molecule. Fig. 7 shows that protein D

proteins. A distinct reaction could be detected at 0.15

and 0.3 yg of the two ign proteins, respectively. Two
same technique could also distinctly be detected at 0.3 yg

(data not shown). In dot blots ign-Fab fragments and ign
Fc fragments bound protein D at 2.5 and 1.2 yg, respectively. In contrast 8 different igc myeloms proteins

representing all subclasses and L-chain types showed no

representing all subclasses and L-chain types showed no

visible reaction with protein D at 5 yg. Neither could any

visible reaction with protein D at 5 yg. Neither could any

visible reaction with protein D at 5 yg. Neither could any

visible reaction with protein D at 5 yg. Neither could any

visible reaction with protein D at 5 yg. Neither could any

visible reaction with protein D at 5 yg. Neither could any

visible reaction with protein D at 5 yg. Neither could any

visible reaction with protein D at 5 yg. Neither could any

visible reaction with protein D at 5 yg. Neither could any

IgG a weak reaction was detected at 5 µg (Fig. 7).

additional proteins be detected. However, with polyclonal

Cloning of the protein D gene

DNA isolated from H. influenzae 772 was partially digested with Sau3A and enriched for fragments in the size of 2 to 7 kilobasepairs (kbp) by fractionation on a 5 sucrose gradient. These fragments were ligated to the BamHI-cut and phosphatase-treated vector pUC18. E. coli JM83 cells transformed with the ligation mixture by high voltage electroporation were plated selecting for resistance to ampicillin. Individual colonies were transferred to nitrocellulose filters and screened with a cocktail of monoclonal antibodies (MAbs) as described in Materials and Methods

Among the 15,000 colonies tested, 60 were found positive. Eight positive colonies were picked, purified and subjected to another two rounds of screening. All clones remained positive during the purification. The purified clones were tested for IgD binding with human IgD, rabbit anti-human IgD and peroxidase conjugated goat anti-rabbit Ig:s in a colony immunoassay as described in Materials and Methods. All were positive regarding IgD binding. Additionally, the clones were found positive when screening with the three MAbs individually.

Restriction_enzyme analysis of plasmid DNA from the positive clones showed that all but one clone carried a 3.3 kbp insert with two internal Sau3A sites. One clone contained an additional 2.0 kbp Sau3A fragment. One of the smaller recombinant plasmids, pHIJ32, was chosen for further characterization. A partial restriction enzyme map was established for the insert of H. influenzae DNA in pHIJ32 (Fig. 8). To identify the region coding for protein D, restriction enzyme fragments were subcloned into pUC18. The resulting transformants were tested for expression of protein D using colony immunoblot analysis as described above. These experiments showed that plasmids carrying a 1.9 kbp HindIII-ClaI fragment-from one end of the insert allowed expression of IgD-binding protein. This recombinant plasmid, called pHIC348, was kept for further experi-

SI

ments. The protein D gene cloned in pHIC348 is expressed from a promoter in pUC18. This was shown by cloning the tion in pUC19. All transformants expressed 1gD binding, as would be expected if the gene is under the control of an endogenous promoter. Transformants carrying the Hindill-Clai fragment in the opposite direction to pHIC348 grew poorly and autolysed during cultivation. This was probably due to the lac2 promoter of pUC19 being oriented in the same direction as the promoter of protein D which led to an overexpression of protein D which was lethal to the cells. In pHIC348 the lac2 promoter was in the opposite direction of the protein D which was in the opposite cells. In pHIC348 the lac2 promoter was in the opposite direction of the protein D promoter was in the opposite

DNA sequence analysis of the protein D gene

The nucleotide sequence of both strands of the insert

from pHIC348 was determined either by direct plasmid sequencing of subclones and deletion constructs or by subcloning restriction fragments into phages Ml3mpl9. Commersially available universal and reverse Ml3mpl9. Setting and the sequencing strategy is outlined in Fig. 8.

The DNA sequence (Fig. 9) reveals an open reading frame of 1092 bp starting with an ATG codon at position 204 and finishing at position 1296 with a TAA stop codon. The open reading frame corresponds to a protein of 364 amino acid residues. Ten nucleotides upstream of the methioning codon is a sequence, AAGGAG, that is complementary to the 3' end of the 165 rRNA of E. coli (Shine, 1974). The spacing between the centre of this putative ribosome-binding site (Tbs) and the start codon is 13 bp 1974). The spacing between the centre of this putative ribosome-binding site (Tbs) and the start codon is 13 bp 1974). The spacing between the centre of this putative ribosome-binding site (Tbs) and the start codon is 13 bp 1974). The spacing site (Tbs) and the start codon is 13 bp 2074). The spacing site (Tbs) and the start codon is 13 bp 2074 and 2074 are spacing site (Tbs) and the start codon is 13 bp 2074 and 2074 are spacing site (Tbs) and the start codon is 13 bp 2074 are spacing site (Tbs) and the sequences of the -10 region, TAAAAT (151-156), and the -35 region, the -10 region, TAAAAT (151-156), and the -35 region,

TTGCTT (127-132), show homology to the consensus of E.

21

coli promoters (Rosenberg, M. and Court, D., Annu. Rev. Genet, 13:319, 1979) and are identical with promoters recognized by the E. coli RNA polymerase. The spacing between the putative -10 and -35 sequences is 18 bp, which is comparable with the favoured value of 17 bp.

Between position 1341 and 1359 there is an inverted repeat with the potential to form a stem and loop structure. This repeat does not, however, resemble a typical rho-independent transcription terminator.

10 Protein D structure

The gene for protein D encodes for a protein of 364 amino acid residues deduced from the nucleotide sequence (Fig. 9). The N-terminal amino acid sequence has typical characteristics of a bacterial lipoprotein signal peptide 15 (Vlasuk et al., J. Biol. Chem. 258:7141, 1983) with its stretch of hydrophilic and basic amino acids at the N-terminus followed by a hydrophobic region of 13 residues, and with a glycin in the hydrophobic core. The putative signal peptide ends with a consensus sequence Leu-Ala-Gly-Cys. 20 recognized by the enzyme signal peptidase II (SpaseII). ~ The primary translation product has a deduced molecular weight of 41,821 daltons. Cleavage by SpaseII would result in a protein of 346 amino acids with a calculated molecular size of 40,068 daltons, in contrast to the 25 estimated size of the mature protein D of approximately 42 kilodaltons. Posttranslational modifications of the preprotein may account for this discrepancy. Several attempts to determine the amino-terminal amino acid sequence of protein D were performed by applying about 1000 pmoles thereof in an automated amino acid sequencer. Since no amino acid phenylthiohydantoin derivatives were obtained, the amino-terminal end of the single IgD-receptor polypeptide chain is probably blocked.

Protein D expressed in E. coli JM83 carrying pHIC348 was analysed in immunoblotting experiments (Fig. 10).

Cytoplasmic, periplasmic and membrane fractions from cells in late logarithmic phase were separated on a SDS-PAGE gel

YAAMMUS

and electroblotted to an Immobilon filter. A protein that binds all three anti-protein D monoclonal antibodies (16C10, 20G6 and 19B4) and radiolabeled IgD could be detected in all three fractions (lane 2-4) from E. coli weight of 42 kilodaltons, i.e. equal or similar to protein D prepared from H. influences (lane 1, Fig. 10).

The nucleotide sequence and the deduced amino acid

sequence of H. influences 772 protein D were compared with using a computer search in the EMBL and Genbank Data using a computer search in the EMBL and Genbank Data nibraries. Apart from similarities in the signal sequence no homology was found.

ren-Ala-Gly-Lys for bacterial lipoproteins. amino acids containing a consensus sequence, 41,821 daltons including a putative signal sequence of 18 sciq sedneuce corresponding to a molecular weight of as well as the nucleotide sequence and the deduced amino protein D gene from H. influenzae in E. coli is described purification of protein D is described. Cloning of the antibodies and monoclonal human igb. A method for strains interacts with three different mouse monoclonal weight immunogenic similarities since protein D from all strains shows in addition to the same apparent molecular isolates of H. influenzae studied. The protein from all detected in all of 116 encapsulated and non-encapsulated apparent molecular weight of 42,000. Protein D can be named protein D is an Ig receptor for human IgD and has an related Haemophilus species is described. The protein A novel surface exposed protein of H. influenzae or SI

32

15

CLAIMS

- A surface exposed protein, which is conserved in
 many strains of Haemophilus influenzae or related Haemophilus species, having an apparent molecular weight of
 42,000 and a capacity of binding human IgD, or naturally occurring or artificially modified variants thereof, or an immunogenic or IgD-binding portion of said protein or
 variants.
 - 2. A protein according to claim 1, having the amino acid sequence as described in Fig. 9, or naturally occurring or artificially modified variants thereof, or an immunogenic or IgD-binding portion of said protein or variants.
- 3. A plasmid or phage containing a genetic code for a protein of Haemophilus influenzae or related Haemophilus species, said protein having an apparent molecular weight of 42,000 and a capacity of binding human IgD, or for naturally occurring or artificially modified variants thereof, or for an immunogenic or IgD-binding portion of said protein or variants.
- A non-human host containing a plasmid or a phage as defined in claim 3 and capable of producing said
 protein or variants or a portion of said protein or variants, which host is chosen among bacteria, yeasts and plants.
 - 5. A host according to claim 4, c h a r a c t e r i s e d in that it is E. coli.
- 30 6. A DNA segment comprising a DNA sequence which codes for a protein of Haemophilus influenzae or related Haemophilus species, said protein having an apparent molecular weight of 42,000 and a capacity of binding human IgD, or for naturally occurring or artifically modified variants thereof, or for an immunogenic or IgD-binding portion of said protein or variants.

7. A DNA-segment according to claim 6, wherein the

8. A recombinant DNA molecule containing a nucleotide DNA sequence is the one specified in Fig. 9.

occurring or artifically modified variants thereof, or for and a capacity of binding human 1gb, or for naturally said protein having an apparent molecular weight of 42,000 Haemophilus influenzae or related Haemophilus species, sequence coding for a surface exposed protein of

variants, which nucleotide sequence is fused to another an immunogenic or IgD-binding portion of said protein or

sequence according to claim 8. 9. A plasmid or phage containing a fused nucleotide

or phage according to claim 9, which host is chosen among 10. A non-human host containing at least one plasmid

11. A host according to claim 10, c h a r a c pacteria, yeasts or plants.

12. A fusion protein or polypeptide in which a surterised in that it is E. coli.

tein by the use of a recombinant DNA molecule according to of said protein or variants, is combined with another provariants thereof, or an immunogenic or IgD-binding portion igb, or naturally occurring or artificially modified cular weight of 42,000 and a capacity of binding human Heemophilus species, said protein having an apparent moleisce exposed protein of Haemophilus influenzae or related

.8 misto

naturally occurring or artificially modified variants of 42,000 and a capacity of binding human IgD, or species, said protein having an apparent molecular weight protein of Haemophilus influenzae or related Haemophilus 13. A fusion product in which a surface exposed

bound to a protein, carbohydrate or matrix. protein or variants, is covalently or by any other means thereof, or an immunogenic or igD-binding portion of said

20

- 14. A vaccine containing a surface exposed protein of Haemophilus influenzae or related Haemophilus species, said protein having an apparent molecular weight of 42,000 and a capacity of binding human IgD, or naturally 5 occurring or artificially modified variants thereof, or an immunogenic or IgD-binding portion thereof.
 - 15. A vaccine containing a surface exposed protein of Haemophilus influenzae or related Haemophilus species, said protein having an apparent molecular weight of 42,000 and a capacity of binding human IgD, or naturally occurring or artificially modified variants thereof, or an immunogenic or IgD-binding portion thereof, combined with another vaccine.
- 16. A vaccine containing a surface exposed protein of 15 Haemophilus influenzae or related Haemophilus species, said protein having an apparent molecular weight of 42,000 and a capacity of binding human IgD, or naturally occurring or artificially modified variants thereof, or an immunogenic or IgD-binding portion thereof, combined with an immunogenic portion of another molecule.
 - 17. A hybridoma cell capable of producing a monoclonal antibody to an immunogenic portion of a surface exposed protein of Haemophilus influenzae or related Haemophilus species, said protein having an apparent molecular weight of 42,000 and a capacity of binding human IgD.
- 18. A purified antibody which is specific to an immunogenic portion of a surface exposed protein of Haemophilus influenzae or related Haemophilus species, 30 said protein having an apparent molecular weight of 42,000 and a capacity of binding human IgD.
- 19. A method of detecting the presence of Haemophilus influenzae or related Haemophilus species in a sample by contacting said sample with the antibody of claim 18 in 35 the presence of an indicator.

tor a surface exposed protein of Haemophilus influenzae or constructed to correspond to the nucleic acids which code contacting said sample with a DNA probe or primer influenzae or related Haemophilus species in a sample by 20. A method of detecting the presence of Haemophilus

human IgD, or for naturally occurring or artificially rent molecular weight of 42,000 and a capacity of binding related Haemophilus species, said protein having an ppa-

-binding portion of said protein or variants. modified variants thereof, or for an immunogenic or IgD-

of 42,000 and a capacity of binding human IgD, or species, said protein having an apparent molecular weight protein of Haemophilus influenzae or related Haemophilus ΣI . A method of detecting IgD using a surface exposed

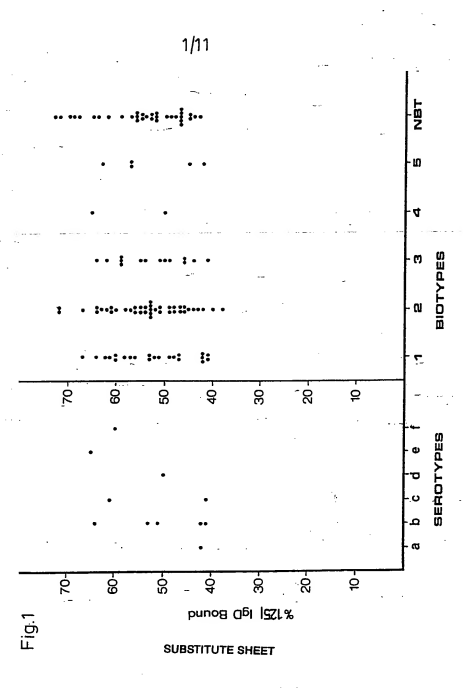
matrix. protein or variants, optionally labelled and/or bound to a thereof, or an immunogenic or IgD-binding portion of said naturally occurring or artificially modified variants

IgD, or naturally occurring or artificially modified molecular weight of 42,000 and a capacity of binding human Haemophilus species, said protein having an apparent exposed protein of Haemophilus influenzae or related 22. A method of separating IgD using a surface

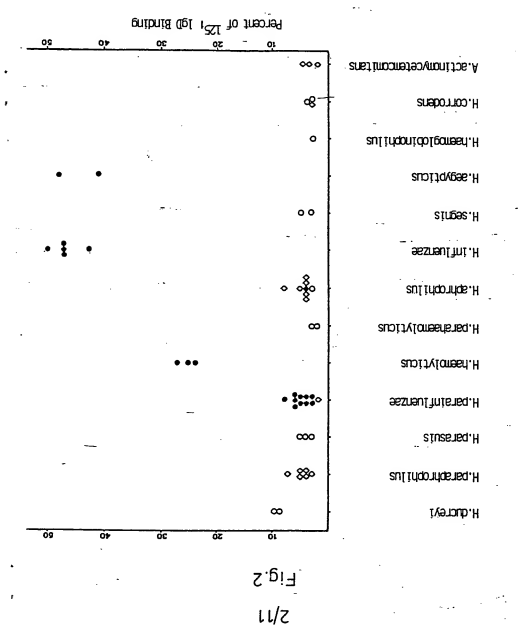
25 of said protein or variants, optionally bound to a matrix. variants thereof, or an immunogenic or IgD-binding portion

30

32



BNSDOCID: <WO__9118926A1_I_



SUBSTITUTE SHEET

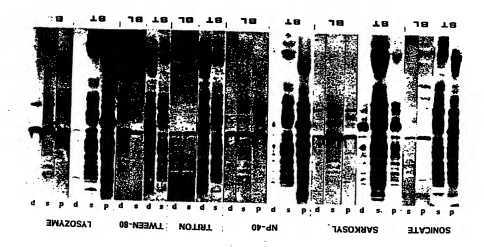
3/11

Fig.3

	STAIN_					BLOT									
			772				772					3198		Minn A	
	Mr	s			SS	_	S	þ	р	SS	р	SS	P	SS	
94 kDa —		=	=		•	•	٠							_	
67 kDa		===			•									:	
43 kDa —	-						_	_	_	_		·	~		
30 kDa		T STATE OF THE STA		_		:								: .·	
20 kDa —	_	**	223				-							-, <i>:</i>	

SUBSTITUTE SHEET

THE SHEET

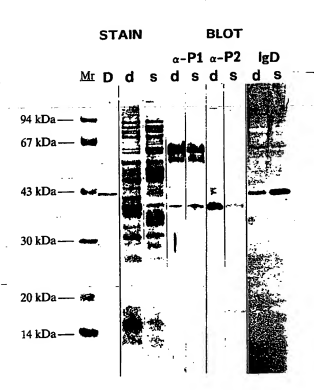


7.6jJ

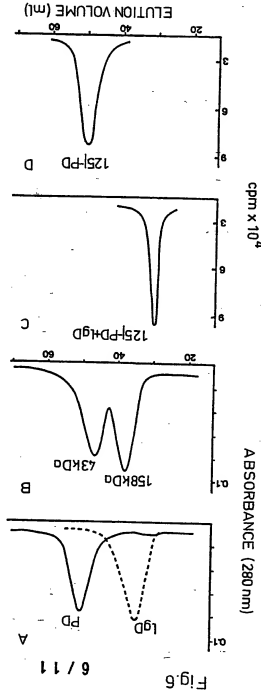
11/7

5/11

Fig.5

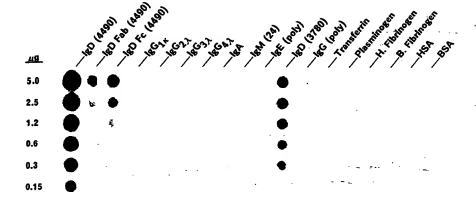


SUBSTITUTE SHEET

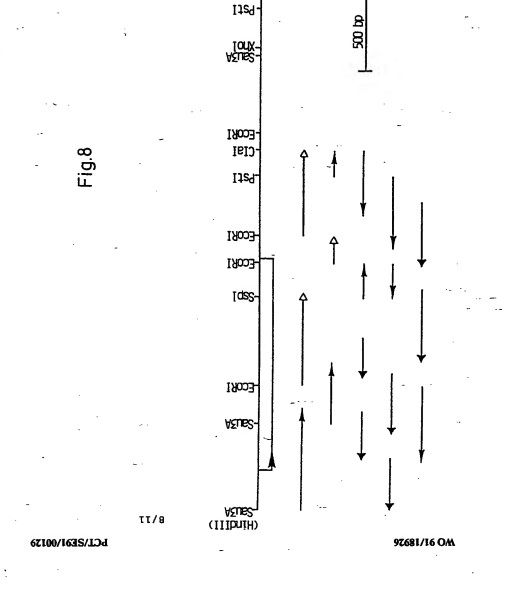


7/11 -

Fig.7



SUBSTITUTE SHEET



ASUBS -

Ind).

9/11

Fig.9a

.08	AAAAAAGGCGGTGGGCAAATTGCTTAGTCGCCTTTTTTGTAACTAAAAATCTAAAAACTCT	167
•	-35 -10	
L6B	ATAAAAATTTACCGCACTCTTAAGGAGAAAATACTTATGAAACTTAAAACTTTAGCCCTT	227
228	TCTTTATTAGCAGCTGGCGTACTAGCAGGTTGTAGCAGCCATTCATCAAATATGGCGAAT SerLeuLeuAlaAlaGlyValLeuAlaGlyCysSerSerHisSerSerAsnMetAlaAsn	287
88	ACCCAAATGAAATCAGACAAAATCATTATTGCTCACCGTGGTGCTAGCGGTTATTTACCA ThrGlnHetLysSerAspLysIleIleIleAlaHisArgGlyAlaSerGlyTyrLeuPro	347
148	GAGCATACGTTAGAATCTAAAGCACTTGCGTTTGCACAACAGGCTGATTATTTAGAGCAA GluHisThrLeuGluSerLysAlaLeuAlaPheAlaGlnGlnAlaAspTyrLeuGluGln	407
801	GATTTAGCAATGACTAAGGATGGTCGTTTAGTGGTTATTCACGATCACTTTTTTAGATGGC AspLeuAlaMetThrLysAspGlyArgLeuValValIleHisAspHisPheLeuAspGly	467
468	TTGACTGATGTTGCGAAAAAATTCCCACATCGTCATCGTAAAGATGGCCGTTACTATGTC LeuThrAspValAlaLysLysPheProHisArgHisArgLysAspGlyArgTyrTyrVal	527
528	ATCGACTTTACCTTAAAAGAAATTCAAAGTTTAGAAATGACAGAAAACTTTGAAACCAAA IleAspPheThrLeuLysGluIleGlnSerLeuGluMetThrGluAsnPheGluThrLys	587
88	GATGGCAAACAAGCGCAAGTTTATCCTAATCGTTTCCCTCTTTGGAAATCACATTTTAGA AspGlyLysGlnAlaGlnValTyrProAsnArgPheProLeuTrpLysSerHisPheArg	647
548	ATTCATACCTTTGAAGATGAAATTGAATTTATCCAAGGCTTAGAAAAATCCACTGGCAAA IleHisThrPheGluAspGluIleGluPheIleGlnGlyLeuGluLysSerThrGlyLys	707
708	AAAGTAGGGATTTATCCAGAAATCAAAGCACCTTGGTTCCACCATCAAAATGGTAAAGAT	767

CURSTITUTE SHEET

โเ/0เ

Ì

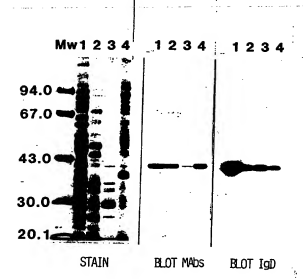
97681/16 OM

49£ [CAACATCAAAATCAAAAATCTAATTTTTAAATTTAAAAAA	8057
2001	TITACTGATITCCCAGATATCTGGCGTGGAATTAAAAGGAATAAAATATATCCCTCA PheThrAspPheProAspThrGlyValGluPheLeuLysGlylleLysEnd	7548
1547	TTCACAGACGTAAATCAAATGTATGCCTTATTGAATAAATCAGGGGCAAGGTGTAA TTCACAGACGTAAAATCAAATGTATGTATATGAAAATCAGGGGCCAACGTGTA PheThraspValAanGtAYTATAATGCCTTATTGAATAAATCAGGGGCCAACGTGTA	7788
. 4811	GCACAATATAATGTGGAAGTGCATCCTTACACCGTGCGTAAAGATGCACTGCCGGAGTTT Alacintytaanvalgluvalhiaptotytthtvalatgiysaapalaleuptogluphe	8ZTT
1753	CTTAATAAGAAGAATCCAAAACCTGATAATATTGTGTACACTCCGTTGGTAAAAGAACTT ValAsniysGluGluSerlysProAspAsnileValTyrThrProLeuValLysGluLeu	1068
190 T	CGTGCAATGGCAGAAGTGGTTAAAATATGCCGAATGGTTGGCCCAGGTTGGTATATGTTA G1yAlamelalagiuvalvaliystytagccgatggtggtggtggtggtgg G1yAlamelalagiuvalvaltatatatata	8007
L 00T	CYYCYYYYYCYCCCYYYCCCLTYLLCCCLYYYCLYLYYYLLYCCYLLCGYLCLLLYYYYCCL CYYCYYYYYYCCCYYYCCLLYLLLCCCLYYYCLYLYYYCCYLLCCYLCLLLYYYYCCL	876
L 76	CAAATGGGAATGGAATTAGTTCAATTAATTGCTTATACAGATTGGAAAGAAA	888
488	CILLYCILYCAYYCLLLCCYLLLLYYYCYYLLYYYCCLYLCYYYCCCYYLCGYFLCILCCY Asjlyrleucharyneyspheysucharyngyrahicharyyccoryntchar	828
758	ATTGCTGCTGAAAACGCTCAAAAAAAAAAAAAAAAAAAA	894

PCT/SE91/00129

11/11

Fig.10



- 1. H.influenzae
- 2. E.coli pHIC348 (cytoplasmic fraction) -
- 3. E.coli pHIC348 (periplasmic fraction)
- 4. E.coli pHIC348 (membrane fraction)

SUBSTITUTE SHEET

INTERNATIONAL SEARCH REPORT

COLD PCT/126/210 (Second apest) (January 1885)
COLD PCT/126/210 (Second apest) (January 1885) Mikael Berustrand/Yvonne Sibsteen unach. TECHTO PERIODINA TO STUTENDIS International Searching Authority 8 2 -90- 1661 1661 թոսն վե**հ**Տ Date of Mailing of this International Search Report Date of the Actual Completion of the International Search IV. CERTIFICATION "a" document member of the same patent family document referring to an oral disclosure, use, exhibition or other means "L" document which may throw doubts on mindiy cisim(s) or which is clied to establish the publication date of secolest citation or other special reason (se specified) "F" earlier document but published on or after the data and contextings and class and space applications of contexting or the contexting of "A" decoment defining the general state of the art which is not considered to be of particular relevance * Special categories of cited documents: 10 see the whole document 25 October 1989, I-SS AZ, 0338265 (AMERICAN CYANAMID COMPANY) see page 3379 - page 3384 "del nemuH rolytinillA diw The Journal of Immunology, Vol. 145, November 1990 Macrong Ruan et al.: "Protein D of Haemophilus influenzae A Novel Bacterial Surface Protein I-22 X'd see bage 119 - page 125 influenzae: Cloning, Aucleotide Sequence, and Expression in Escherichia coli^u, zulindomesH To mistory Profing Profesion of Haemophilus 1991 Håkan Janson et al: "Protein D, an Infection and Immunity, Vol. 59, No. 1, January 1-22 X'd Citation of Document, 11 with indication, where appropriate, of the relevant passages 12 Relevant to Claim Ho.13 III. DOCUMENTS CONSIDERED TO BE RELEVANT SE, DK, FI, NO classes as above Documentation Searched other than Minimum Documentation to the Extent that such Documents are Included in Fields Searched⁸ C 07 K; C 12 N; A 61 K **Ibc2** Classification Symbols Classification System Minimum Documentation Searched IF LIETDS SEVECHED C 15 N 12/62 89/1 0 21 3 IPCS: C 07 K 15/04, C 12 N 15/31, A 61 K 39/102//C 12 Q 1/04 According to International Patent Classification (IPC) or to both Mational Classification and IPC L CLASSIFICATION OF SUBJECT MATTER (it soverel classification symbols apply, indicate ail) International Application No PCT/SE 91/00129

III. DOCI Category	MENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET) Citation of Occument, with indication, where appropriate, of the relevant passages	Relevant to Claim No
A	The Journal of Immun logy, Vol. 122, No. 4, April 1979 Arne Forsgren et al.: "Many bacterial species bind human IgD", see page 1468 - page 1472	1-22
·		
		•
	·	
	·	
	· ·	
ĺ		
	· ·	
	- 142 - 14	
	• 1	
		•
		. •
	;	
	<u></u>	
ì	,	

CM INTERNATIONAL PATENT APPLICATION NO.PCT/SE 91/00129

This snex lists the patent samily members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the Swedish Patent Office EDV files on The Swedish Patent Office is in no way lishle for these particulars which are merely given for the purpose of information.

edab	(s)rada	79W	gleb	Patent document cited in search report
86-10-56 86-10-58	987£1££ 4283702	-G-UA -A-q€	97-01-68	EP-A2- 0338265
		·		-
-				
			·	
•	•			
				·
			:	
	· -			
Ÿ.				
			-	
	•			_ .
•	**.			•